Features of vaginal microbiocenosis in women of reproductive age with overweight and obesity

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Introduction

Obesity is a significant medical and social problem of our time, the relevance of which is primarily related to the growing prevalence and the threat of the formation of various pathological human conditions [1, 2, 18, 23, 27]. Obesity and overweight play an important negative role in gynecological and obstetric practice. There is information...
that in women against the background of metabolic disorders increases the frequency of infectious pathology of various habitats, including the urogenital tract [6, 12, 19, 20, 28].

According to the prevalence of urogenital infections in women, one of the leading places is urogenital candidiasis [3, 21, 22, 26]. The most common causative agent of candidiasis is the fungus Candida albicans. But in the last decade in the etiological structure of candidiasis there is a tendency to increase the proportion of fungi Candida non-albicans: C. glabrata, C. tropicalis, C. parapsilosis, C. krusei [10]. A sign of modern genital candidiasis is the formation of associations of fungi of the Candida with various representatives of opportunistic pathogen S. Associations with Candida fungi may include gram-positive and gram-negative aerobic, facultative-anaerobic and obligate-anaerobic microorganisms [29].

Firmicutes have a significant proportion in the bacterial spectrum of the vagina. Catalase-positive and coagulase-negative staphylococci in the vaginal microbiocenosis occur in more than 50% of women, Streptococcus spp. and Enterococcus spp. - in 30-40% of case S. Anaerobic gram-positive cocci (Peptococcus spp., Peptostreptococcus spp.) are detected with a frequency of 30-80%. Gram-negative anaerobic rod-shaped bacteria contaminate the female genital tract with considerable frequency. Prevotella spp., Bacteroides spp., Porphyromonas spp., Fusobacterium spp., Veillonella spp [17]. Among enterobacteria, as factors of infective pathology of genitals, a significant role belongs to Escherichia coli, Klebsiella spp., Enterobacter spp. and others [13]. Lactobacilli occupy a dominant position in the microflora of the vagina and cervix. Bifidobacterium are excreted from the genital tract with less frequency. Lactobacilli are a marker of the normal microbiocenosis of the genital tract. It is lactobacilli that determine the degree of colonization resistance of the vagina [8, 9, 15, 16].

One of the most common dysbiotic disorders of the vaginal microbiota is bacterial vaginosis (BV), which should be considered as an infectious non-inflammatory syndrome associated with dysbiosis of the vaginal habitat [5, 24, 25, 30].

The main manifestations of this pathology are a decrease or absence in the microflora of the vagina lactobacilli and activation of Atopobium vaginae and Gardnerella vaginalis, which play a "key" role in the pathogenesis of bacterial vaginosis. Atopobium vaginae is able to block factors of nonspecific reactivity (complement system), leads to increased inflammation in bacterial vaginosis, is a competitor of lactobacilli in terms of glycogen consumption. Data from the scientific literature indicate that metabolic disorders in women contribute to the active growth of Gardnerella vaginalis, which produces a specific toxin (cytolysin) that can cause desquamation of vaginal endothelial cells [4, 7, 14].

An important component of the bacterial spectrum in bacterial vaginosis is Mobiluncus spp. This microorganism produces a toxin that inhibits the adhesion of lactobacilli on vaginal epithelial cells [11].

Given the high prevalence at the present stage of candidiasis and bacterial vaginosis of the genital tract, the negative impact of these diseases on the female reproductive system, the aim of our work was to study changes in vaginal microbiome in patients of these categories to improve existing treatment regimens.

Materials and methods

In order to solve the tasks of our work, we surveyed 120 women of reproductive age with overweight and obesity. For comparison, 60 women without overweight and obesity were examined. All women were divided into groups: Group I - women with vulvovaginal candidiasis and overweight and obesity (60 women); Group II - women with vulvovaginal candidiasis without excess weight and obesity (30 women); Group III - women with bacterial vaginosis and overweight and obesity (60 women); Group IV - women with bacterial vaginosis without excess weight and obesity (30 women). Data obtained from a survey of 30 healthy women were used as controls. Carrying out of microbiological researches and the account of results was carried out according to the Order №234 of the Ministry of Health of Ukraine from 10.05.2007 and other regulatory documents.

The following differential diagnostic media were used to inoculate the genital mucus: blood agar, yellow-salt agar, Endo, Sabouraud agar, MRS medium for lactobacilli, Blaurok medium for bifidobacteria. Crops were carried out by the method of sector seeding on dense nutrient media, which allows to determine the degree of microbial contamination and to identify the maximum possible spectrum of aerobic and facultative anaerobic microflora.

Gardnerellosis was diagnosed by bacterioscopy by staining with Romanovsky smears with subsequent counting of "key" cells, amine test, determination of pH. The contents of the vagina of women for the presence of fungi Candida were examined by microscopy and culture bacteriological method. Sabouraud agar was used to identify yeast-like fungi. MICROMA-TEST kits "Candidatest 21" (Erba Lachema s.r.o. (Czech Republic)) were used for further identification of isolated yeast-like fungi.

Determination of anaerobic microorganisms was carried out according to the guidelines "Laboratory diagnosis of purulent-inflammatory diseases caused by asporogenic anaerobic microorganisms", Kharkiv, 1985. The anaerobic microflora of the genital tract was studied with strict adherence to the technique of anaerobic cultivation. Thioglycollate medium, blood agar with glucose, liver broth, Kitt-Tarozzi medium were used for the study. The "Anaerocult" system (Merck, Germany) and the "Anaerogas" gas packages were used to create anaerobic conditionS. MICROMA-TEST "Anaerotest 23" kits (Erba Lachema s.r.o. (Czech Republic)) were used for further identification of isolated anaerobic microorganisms.
Identification of microorganisms was performed by their cultural and morphological characteristics. When identifying the isolated microorganisms, the Bergey's classification was followed.

To assess the contamination of the genital tract was carried out quantitative accounting of colonies that grew in dense nutrient media. The content of opportunistic pathogens in the test material was expressed by the number of colony-forming units in 1 ml (CFU/ml) of biological material.

Statistical analysis of the obtained research results was performed using standard computer packages "Data Analysis" Microsoft Excel for Windows 2002.

**Results**

The results of microscopy of the contents of the vagina in most women of group I with vulvovaginal candidiasis indicate negative changes in the microbiota, which have manifestations of vaginitis. In 83.3% of patients there was a significant increase in the number of neutrophils and leukocytes (30-100 in the field of view), increased epithelial desquamation, degenerative changes of the nuclei, vacuolation of the cytoplasm, deficiency or absence of normal morphotype microflora, a significant increase in gram-positive and gram-negative organisms. In 46.7% of the examined fungi were in the form of yeast cells, in 53.3% yeast and pseudomycellium were found.

In 6.7% of obese women, against the background of a moderate number of yeast cells, no manifestations of vaginitis were detected, but there was an accumulation of gram-positive cocci and gram-negative rods, as well as an increase in root-like bacteria and fusobacteria.

Examination of the contents of the vagina by bacteriological culture revealed that in patients of group I, Candida fungi were sown in all subjects at a significant concentration (lg5.8 CFU/ml).

In 95% of patients the formation of associations of fungi of the Candida with various representatives of opportunistic pathogenic microflora was revealed. Two-component bacterial-fungal associations were found in 28.3% of cases, three-component - in 48.3%, four-component - in 18.3% of patients. Monocultures of Candida fungi were registered in 5.1% of women.

Associative forms of vaginal dysbiosis differ from monomicrobial ones by greater aggressiveness, significant resistance to antibacterial drugs, severe structural changes in the vaginal mucosa.

As a result of bacteriological examination of women of group I, an increase in the composition of bacterial-fungal associations of the vaginal contents of Firmicutes, in particular, cocci with pathogenic properties was found. Thus, the frequency of the presence of S.aureus in the associations in women of group I reached 18.3%, epidermal staphylococcus with hemolysis - 23.3%, enterococcus - 20.0%.

Also, the associations of women of group I with a significant frequency included different types of enterobacteria. A significant proportion was found in the associations of Escherichia coli with hemolytic properties (16.7%), Klebsiella spp. - 15%, Enterobacter spp. - 13.3%. Quantitative seeding rates of various representatives of opportunistic vaginal microflora in patients exceeded the diagnostic level (lg4.2 - lg5.8 CFU/ml) (Fig. 1).

E. coli, E.coli, S.pyogenes, S.agalactiae, E.faecalis, S.aureus, S.epidermidis (hern), S.epidermidis

![Fig. 1. Comparative diagrams of the composition of vagina microflora healthy women and women of reproductive age with vulvovaginal candidiasis and obesity.](image1)

Lactobacillus deficiency was found in 58.3% of patients, and their absence - in 10.0%. In 31.7% of cases, the registration rates of lactoflora were normal. Also in women of group I there was a decrease in the quantitative level of lactobacilli.
seeding lactobacilli.

According to the results obtained with the highest frequency, the vagina of patients of group I was contaminated with Peptostreptococcus spp. (28.3%), Fusobacterium spp. (16.7%) and Veillonella spp. (16.7%). The presence of Bacteroides spp. (13.3%), Prevotella spp. (10.0%), Clostridium spp. (6.7%) were registered with less frequency. The frequency of registration of Gardnerella vaginalis was only 3.3%, Mobiluncus spp. - 1.7%.

Quantitative seeding rates of individual members of the obligate anaerobic microflora exceeded the diagnostic level: Peptostreptococcus spp. - lg5.2 CFU/ml, Veillonella spp. - lg5.4 CFU/ml, Fusobacterium spp. - lg4.4 CFU/ml.

Quantitative level of vaginal contamination by Bacteroides spp., Clostridium spp., Prevotella spp. was reduced compared to normal (lg2.0 - lg2.8 CFU/ml).

In order to compare the indicators of vaginal microecology, the second group of patients with vulvovaginal candidiasis without obesity was examined.

Examination of the contents of the vagina in patients of group II by microscopy shows a less significant than in patients of group I, an increase in the number of leukocytes (15-50 in the field of view), moderate destructive changes in epithelial cells, lower frequency of pseudomycelial form of fungi (yeast cells - 76.7%, yeast cells and pseudomyecelium - 33.3%). Microorganisms belonging to lactobacilli by morphotype were found more frequently in obese women than in group I patients.

According to the results of the study of vaginal contents by bacteriological method in women of the second group, despite the significant frequency of formation of fungal-bacterial associations (83.3%), the quantitative indicators of individual microbiocenosis in the subjects did not reach the level found in patients of group I. Two-component associations were found in 46.7% of the surveyed, three-component associations in 36.7% and monocultures in 16.6%.

The concentration of Candida fungi exceeded the norm and was lg4.6 CFU/ml. Gram-positive coccal microflora with pathogenic properties and enterobacteria were associated with fungi with a lower frequency than in patients of group I, and in low concentration. In women of group II, the frequency of vaginal contamination of S.aureus was 13.3%, S.epidermidis (hem+) - 16.7%, Enterococcus faecalis - 16.7%, S.pyogenes - 6.7%. The frequency of vaginal inoculation with enterobacteria in women was in the range of 10-16.7%. Quantitative indicators of Firmicutes with pathogenic properties exceeded the diagnostic level (>lg4.0 CFU/ml).

Some representatives of the anaerobic microflora in the vaginal associations in women of group II without obesity were registered with a higher frequency than in women of group I: Bacteroides spp. - 36.7%, Clostridium spp. - 10%, Prevotella spp. - 33.3%. The frequency of registration of Gardnerella vaginals was 6.7%, Mobiluncus spp. - 3.3%, Atopobium spp. - 3.3%. Quantitative indicators of anaerobic seeding were in the range of lg3.6 - lg5.0 CFU/ml.

The rates of vaginal contamination with lactobacilli in women of group II were higher compared to women of group I. Lactobacillus deficiency was found in 46.7% of patients.

Quantitative indicators of anaerobic seeding were in the range of lg3.6 - lg5.0 CFU/ml.

Some representatives of the anaerobic microflora in the vaginal associations in women of group II without obesity were registered with a higher frequency than in women of group I: Bacteroides spp. - 36.7%, Clostridium spp. - 10%, Prevotella spp. - 33.3%. The frequency of registration of Gardnerella vaginals was 6.7%, Mobiluncus spp. - 3.3%, Atopobium spp. - 3.3%. Quantitative indicators of anaerobic seeding were in the range of lg3.6 - lg5.0 CFU/ml.

The rates of vaginal contamination with lactobacilli in women of group II were higher compared to women of group I. Lactobacillus deficiency was found in 46.7% of patients.
(6.7%) were detected with the highest frequency, and their quantitative indicators exceeded the diagnostic level. Candida parapsilosis and Candida krusei were sown with insignificant frequency (5% and 3.3%, respectively) (Fig. 2).

The species composition of Candida non-albicans in patients of the second group did not differ much from the species identified in women of group I. However, in the examination of women with vulvovaginal candidiasis without obesity, a lower frequency of vaginal inoculation with Candida non-albicans was found than in patients of group I. Candida tropicalis were detected with the highest frequency (10%) and at a concentration of 4.7 CFU/ml in patients of group II. Other species of non-albicans in women of group II were registered with less frequency and in low concentration. Candida krusei was not detected in the spectrum of non-albicans fungi isolated from the vagina in patients of group II.

The task also included the study of the peculiarities of changes in the microbiocenosis of the vagina in patients with bacterial vaginosis and obesity (group III). As a comparison, patients with vaginosis without obesity (group IV) were examined.

The results of microscopy of the contents of the vagina in women of group III with vaginosis indicate a significant presence of parabasal epithelial cells and "key" cells, the phenomenon of desquamation of the squamous epithelium, a significant density of coccal microflora and rodS. Leukocytes in the vaginal material were registered in moderation (15-35 in the field of view). Morphotypes of microflora corresponding to lactobacilli were detected with insignificant frequency or were absent. The frequency of registration of fungi by microscopy did not reach a high level (yeast cells - 15%, yeast + pseudomycelium - 8.3%).

Bacteriological examination of the contents of the vagina in patients of group III with vaginosis revealed a significant decrease in the frequency and quantitative indicators in the spectrum of vaginal microbiota of Bacteroidetes phylum, which includes Bacteroides and Prevotellaceae family and increased vaginal microflora representatives of Firmicutes.

Associations of anaerobic and facultative-anaerobic microflora with dominance of anaerobes were revealed in all examined patients of the III group. Three-component associations of bacterial flora were observed in 38.3% of patients, four-component associations in 45%, and five-component associations in 16.7%.

Examination of the contents of the vagina showed that with the highest frequency in patients of group III were found Gardnerella vaginalis (83.3%), Atopobium vaginae (68.3%), Mobiluncus spp. (41.7%). The presence of Atopobium vaginae in the spectrum of vaginal microflora is a sign of recurrent disease. Also in the composition of bacterial associations in patients of group III, an increase in the frequency of registration of Peptostreptococcus spp. (43.3%), Fusobacterium spp. (36.7%), Veillonella spp. (31.7%). Bacteroides spp., Prevotella spp. and Clostridium spp. were determined in associations with insignificant frequency (15.0%, 11.7% and 8.3%, respectively).

Quantitative indicators of vaginal contamination by obligate anaerobes in patients of group III significantly exceeded the norm. The diagnostic level of Peptostreptococcus spp., Fusobacterium spp., Veillonella spp., Atopobium vaginae exceeded the diagnostic level. Whereas quantitative seeding rates of Bacteroides spp. were at a low level - lg3.0 CFU/ml (Fig. 3).

Among gram-negative facultative anaerobic rods in the associations of women with vaginosis with the highest frequency were E.coli (18.3%), E.coli hem+ (13.3%), Klebsiella spp. (16.7%). Quantitative indicators of Firmicutes group III patients and intestinal microflora isolated from the vagina exceeded the diagnostic level (lg4.0 - lg5.0 CFU/ml).

In 16.7% of the surveyed women of group III, the presence of Candida fungi was detected in insignificant concentration (lg3.8 CFU/ml). The spectrum of Candida fungi included C.albicans - 8.3%, C.glabrata - 5.0%, C.tropicalis - 1.7%, C.krusei - 1.7% (Fig. 4).

Diysbiotic changes of the vaginal microbiota in patients of group III were characterized by a significant decrease in vaginal contamination by representatives of the normal microflora - lactobacilli. The seeding rate of lactobacilli was 56.7%, absence - 43.3%. Quantitative seeding rates of lactobacilli were low (lg2.2 CFU/ml).

For comparison, we examined patients with vaginosis without obesity (group IV). According to microscopy, no significant leukocyte reaction was detected in the vaginal material (5-25 in sight). There was a significant increase in the number of "key" cells and an increase in the density of vaginal contamination with gram-positive and gram-negative microflora.

Bacteriological examination of the vaginal microbiome in patients with obesity without vaginosis revealed a lower frequency and lower quantitative level of vaginal contamination with obligate anaerobes and facultative anaerobic microflora. The anaerobic spectrum of the vaginal microflora included Gardnerella vaginalis (73.3%), Atopobium vaginae (43.3%), Mobiluncus spp. (30.0%), Peptostreptococcus spp. (33.3%) and Fusobacterium spp. (20.0%). The frequency of registration of anaerobes exceeded that found in patients of group III. Bacteroides spp. - 56.7%, Veillonella spp. - 46.7%, Clostridium spp. - 46.7%.

Quantitative indicators of obligate anaerobic microflora isolated from the vagina of women of group IV were in the range of lg4.4 - lg6.0 CFU/ml. But in women of group IV, the seeding rates of Bacteroides spp. and Prevotella spp. significantly exceeded those found in patients of group III (respectively lg6.0 - lg5.2 CFU/ml).

With a lower frequency than in patients of group III, in patients of group IV, the microbial associations included individual representatives of the intestinal microflora: E.coli...
The frequency of seeding from the vagina of Firmicutes patients, in particular, gram-positive coccal microflora did not reach significant values: S.aureus - 10.0%, S.epidermidis (hem+) - 13.3%. Quantitative indicators of vaginal seeding of facultative-anaerobic microflora were in the range of lg3.5 - lg4.8 CFU/ml.

The contents of the vagina in women of group IV with a low frequency (16.7%) and a small amount included Candida fungi (lg2.8 CFU/ml). Candida albicans was detected with the highest frequency (6.7%). The frequency of registration of fungi Candida non-albicans was: C.tropicalis - 3.3%, C.parapsilosis - 3.3% (Fig. 4).

In patients of group IV, the indicators of vaginal contamination with lactobacilli did not reach the norm, but were at a higher level (76.7%) than in patients of group III in the amount of lg4.2 CFU/ml.

The composition of associations in patients of group IV was represented by 2-4 types of microflora: two-component associations (16.7%), three-component (50%), four-component (33.3%). Five-component bacterial associations were not determined.

Discussion

Clinical assessment of the condition of the lower genitalia without taking into account the data of microscopy is subjective and does not always allow to detect vaginal dysbiosis in patients [29].

The results of examination of women with vaginosis (groups III and IV) show that obese patients form multicomponent anaerobic-aerobic bacterial communities with a significant content of Gardnerella vaginalis, Atopobium vaginae, as well as Firmicutes with pathogenic properties on the background of low or no normal microflora. As S.I. Klimnyuk and co-authors pointed out in their study (2019): not only Gardnerella vaginalis but also anaerobes Mobiluncus spp., Bacteroides spp., Mycoplasma hominis and other microorganisms are involved in the development of vaginal dysbiosis, and the only cause of bacterial vaginosis is a violation of the quantitative composition of microorganisms in the vaginal environment on the background of other dysbiotic processes in the body. Complications associated with bacterial vaginosis include chronic inflammation of the internal genitals, the development of adhesions of the pelvic organs, infertility [17].

Some studies suggest that a higher body mass index is associated with increased susceptibility to bacterial vaginosis, but the results are conflicting. Prior to his open cohort study conducted in Mombasa, Kenya, Lookken E.M. with co-authors in 2019 [19] selected women sex workers aged 16 to 45 years. Women who were obese, the authors concluded, had an almost 20% lower risk of developing bacterial vaginosis compared to women with a normal body mass index. The authors continue to suggest that it is appropriate to study the potential mechanisms of this effect, including the possible effects of diet related to obesity changes in the intestinal microbiome and systemic estrogen levels [19].

At the same time, R.T.Brookheart and co-authors in 2019 [6] showed that bacterial vaginosis is one of the most common vaginal diseases in the United State. Recent studies have shown that in obese women, the abnormal microbiota resembles bacterial vaginosis; however, few studies have examined the prevalence of bacterial vaginosis in overweight and obese population. Moreover, it is not known whether there are racial differences in the ratio of obesity to bacterial vaginosis. In their study, the authors examined the correlation between body mass index and bacterial vaginosis, as determined by the Nugent scale, and identified the influence of race. Overweight and obese women were found to have a higher incidence of bacterial vaginosis than lean women, even after adjusting for race-related variables. Among white women, the prevalence of bacterial vaginosis was higher among overweight women and white women with class I and II/III obesity compared to lean white women, a phenomenon not observed among black women. The authors concluded that overweight and obese women had higher Nugent scores and higher levels of bacterial vaginosis compared to lean women. Interesting was the fact that black women had a higher prevalence of bacterial vaginosis, regardless of their body mass index, compared to white women [6].

Obesity is associated with an increased level of susceptibility to infections, however, the results of clinical studies assessing body mass index and vaginitis are contradictory, G.Ventolini and co-authors pointed out in 2017. They found an association between obesity and recurrent vulvovaginal bacterial infections in women of childbearing potential. The authors proved that obesity can be an independent risk factor for vulvovaginal bacterial infections in women of reproductive age due to altered vaginal immunity [28].

In general, the results of the study of vaginal microbiota in obese women allowed us to identify a significant frequency of two variants of microbiocenosis: candidal vaginitis and vaginal vaginosis, which is confirmed by other studies [21, 27]. The identified variants of vaginal microbiome disorders in patients of our study groups have differences in species and quantitative composition of microbial associations, different indicators of vaginal contamination by Candida fungi, different degrees of imbalance between potentially pathogenic and normal microflora, as indicated by other researchers [24, 28].

The obtained data indicate the feasibility of further development of pathogenetically sound schemes for the treatment of disorders of the vaginal microbiocenosis in obese women.

Conclusions

1. The composition of vagina microbiota in women of reproductive age with overweight and obesity (groups I and
features of vaginal microbiocenosis in women of reproductive age with overweight and obesity

II) differs from the composition of the microbiota in women of reproductive age without obesity (groups III and IV) by statistically significant decrease in vaginal contamination by anaerobic microflora, which has a significant metabolic potential: Bacteroides spp. and Prevotella spp. Gram-positive anaerobic and facultative-anaerobic microflora of Firmicutes have a significant share in the spectrum of vaginal microflora in overweight and obese patients, in contrast to non-obese women of reproductive age.

2. In women of reproductive age with vulvovaginal candidiasis who are obese (group I), in contrast to non-obese patients (group II), a higher frequency of fungal-bacterial associations, a higher content in the associations of Firmicutes with pathogenic properties and Proteobacterium, a higher quantitative level of vaginal contamination by the fungi Candida albicans and non-albicans on the background of deficiency or lack of lactoflora observed.

3. The state of microecology in the vagina in women of reproductive age with bacterial vaginosis, overweight and obesity (group III) is characterized by a significant frequency of multicomponent associations, a significant imbalance between the associates of obligate-anaerobic microflora. An increase in vaginosis-associated microflora (Atopobium vaginae, Gardnerella vaginalis, Mobiluncus spp.) and a significant deficiency of Bacteroides spp., Prevotella spp., Lactobacillus spp. was established.

References


ОБІОЦЕНОЗ

ДЛИШКОВ А

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Ключові слова:

albicans на фоні дефіциту або загальної відсутності лактофлори.

кандидозним вульвовагінітом та ожирінням, на відміну від пацієнток без ожиріння, встановлена більш висока частота

Встановлений низький рівень висіву лактобацил (lg2,2 КУО/мл). Таким чином, значну питому вагу в спектр мікрофлори

піхви жінок з вагінозом та ожирінням виявило значні дисбіотичні порушення складу піхвової мікрофлори, трьох-, чотирьох-

трьох- та чотирьохкомпонентні асоціації грибів р.Candida з різними представниками умовно-патогенної мікрофлори.

можливий спектр аеробної та факультативно-анаеробної мікрофлори. У жінок з вульвовагінальним кандидозом, надлишкової

репродуктивного віку з надлишковою вагою та ожирінням. Визначали ступінь мікробного обсімення та виявляли максимально

піхви у жінок з кандидозом та бактеріальним вагінозом з метою удосконалення існуючих схем терапії. Обстежили 120 жінок

vaginae та Gardnerella vaginalis, що відіграють "ключову" роль у патогенезі БВ. Метою роботи стало вивчення змін мікробіому

Peptostreptococcus Mobiluncus sp, Mycoplasma hominis може збільшуватись у 100 разів. Доведено активну Atopobium

вагіни. При бактеріальному вагінозі концентрація анаеробних патогенів Peptostreptococcus sp, Gardner ella vaginalis,

бактеріальні збудники та суттєво зменшується кількість лактобактерій, які зазвичай є у складі бактер іальної флори

з різними представниками умовно патогенної мікрофлори, такими як грампозитивні та грамнегативні аеро бні,


бактеріальний урогенітальний кандидоз, в етіологічній структурі якого визначна роль належить грибам Candida albicans,

частота інфекційної патології на фоні метаболічних розладів. Найбільш розповсюджена форма інфекційно го вагініта - це

Гаспарян К.А., Кондратюк В.К., Пономарьова І.Г., Кондратюк К.О., Дзісь Н.П., Лисяна Т.О.


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